

# LoF update

## Geuvadis RNA Sequencing Project

# Annotation

The variant annotation follows the 1000 Genomes Project Style

Coding annotation is under the VA flag of the INFO field, and contains the following categories:

SNPs:

NONSYNONYMOUS, SYNONYMOUS, PREMATURESTOP, REMOVEDSTOP,  
REMOVEDSTART, SPLICEOVERLAP

Indels:

SPLICEOVERLAP, ENDOVERLAP, STARTOVERLAP, DELETIONNFS, INSERTIONNFS,  
DELETIONNFS, INSERTIONNFS

Transcripts are annotated using Gencode V7 transcript set for consistency. We should discuss updating the transcript set and freezing the version number.

Update from D. Macarthur on IKG LoF annotation

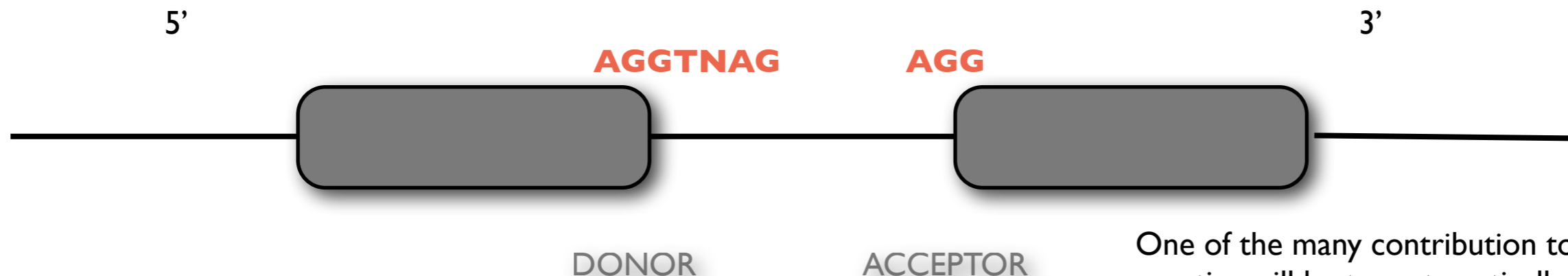
*Described in Geuvadis RNASeq Wiki Page (Tuuli)*  
*22/04/2012*

# Annotation

We propose the following additions/changes for the severe Loss of Function/Protein Truncating variant analysis:

Indels (currently annotated based on positional effect of indel - need more on sequence context work in progress):

1. DONOR\_IN2 - Splice Donor Variants
2. DONOR\_IN\_45AG - Splice Donor Variants
3. DONOR\_EX2\_AG - Exonic Splice Variants
4. ACCEPTOR\_IN2 - Splice Acceptor Variants
5. ACCEPTOR\_EX1\_G - Exonic Acceptor Splice Variants



One of the many contribution to the field of medical genetics will be to systematically assess the relevance of predicted annotated splice variants: Exon Skipping, Intron Retention, etc?

sQTLs will add additional categories that were probably excluded from analysis.

*Additions discussed by Manuel Rivas and Tuuli L.*

*Categories follow Faustino et al. (2003) : Pre-mRNA splicing and human disease, and relevant to human disease: Jordan et al. (2012): PSOR2 is due to mutations in CARD14.*

# Annotation

## Additional meta-information

1. worst=FLAG
2. nmd=BOOL (0 or 1) For frameshift indels and nonsense variants
3. ofptv=BOOL (0 or 1) For splice variants assuming exon-skipping maintain frame?
4. fsX=pepnumber (int) How many downstream peptides for next Stop.
5. pepsize=orig\_new (string) For frameshift indels, nonsense, startlost, and readthrough variants: original peptide length, new peptide length.
6. exin=INT For splice variants: Nearest exon number in transcript. e.g. IVS11+1G>C
7. HGVS=string e.g. c.IVS11+1G>C

*Additions discussed by Manuel Rivas and Tuuli L.*

*Implemented in v0.9 version of PLINK/SEQ software.*

<http://atgu.mgh.harvard.edu/plinkseq/>

# Annotation

LoF/PTV variant	Effect on Transcription	Detection
Nonsense	Nonsense Mediated Decay (NMD)	ASE, Exon Quantification, Transcript Quantification
Splice	NMD, Exon Skipping, Intron Retention	ASE (for exonic splice variants), Splice junction quantification, Exon + Intron Quantification, Transcript Quantification
Frameshift Indels	NMD	Transcript Quantification, Exon Quantification
SV/CNV	Gene Dosage Compensation, eQTL	Transcript Quantification, Exon Quantification
Start Lost	ASE/eQTL	ASE, Transcript Quantification, Exon Quantification
Readthrough	ASE/eQTL	ASE, Transcript Quantification, Exon Quantification

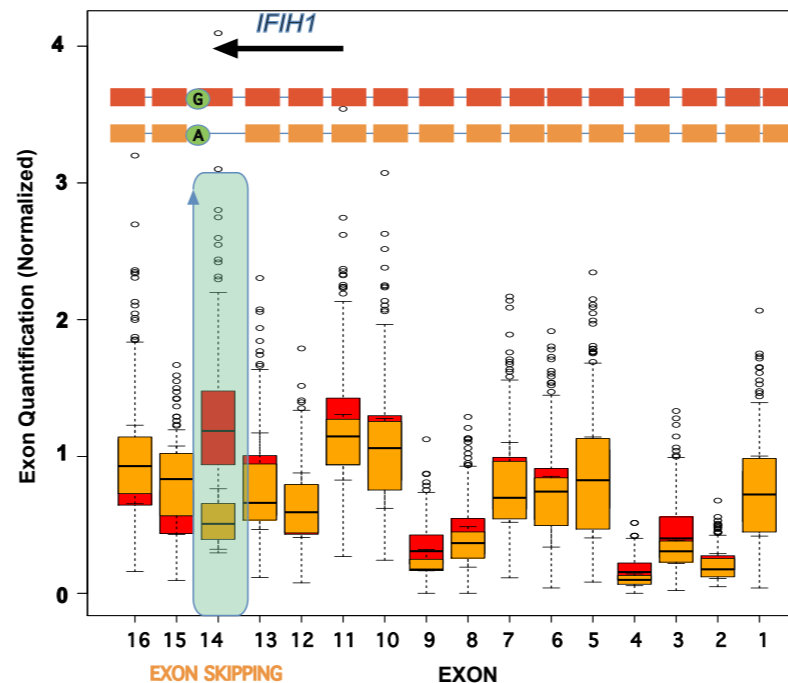
# Quantifications

## I. Exon Quantification

Early data upload made available by Tuuli with BWA mappings.

i. Quantifications from UNIGE are raw counts of reads over exons, calculated by in-house script made by Tuuli.

- 1 we take only uniquely mapped properly paired reads with bwa MAPQ>10 for both mates
- 2 we count reads only in protein-coding and linc-RNA transcripts of the annotation, because we're sequencing a poly-A library
- 3 what is an "exon": we merge all overlapping exons of a gene into a meta-exon, ID: ENSG000001\_exonstartcoord\_exonendcoord. This is to avoid problems with reads that map to several exons that overlap. Note though in the case of partially overlapping exons, our quantification units are not real exons but kind of meta-exons.
4. we count reads over these exons without using information of read pairing, except that we exclude reads where the pairs map to two different genes. We count a read in an exon if either it's start or end coordinate overlaps an exon.
5. in the case of split reads, we count the exon overlap of each split fragment, and add counts per read as  $1/(\text{number\_of\_overlapping\_exons\_per\_gene})$ . I.e. if a read is split and the two parts map to 2 different exons of a gene, we count it as 0.5 in each.



Early p.o.p. of dataset interpretation demonstrates that exon-quantifications are reliable.

*Taken from Geuvadis RNA-Seq wiki page (Tuuli).*

# Methods/Analysis

Early analysis demonstrate strong ASE signal for nonsense variants.

Propose the following additional analyses:

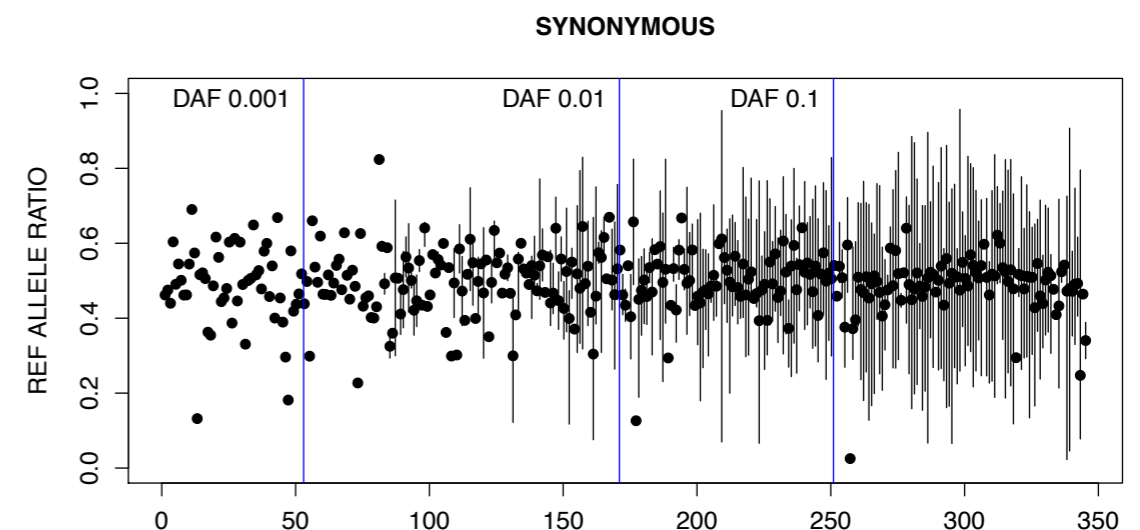
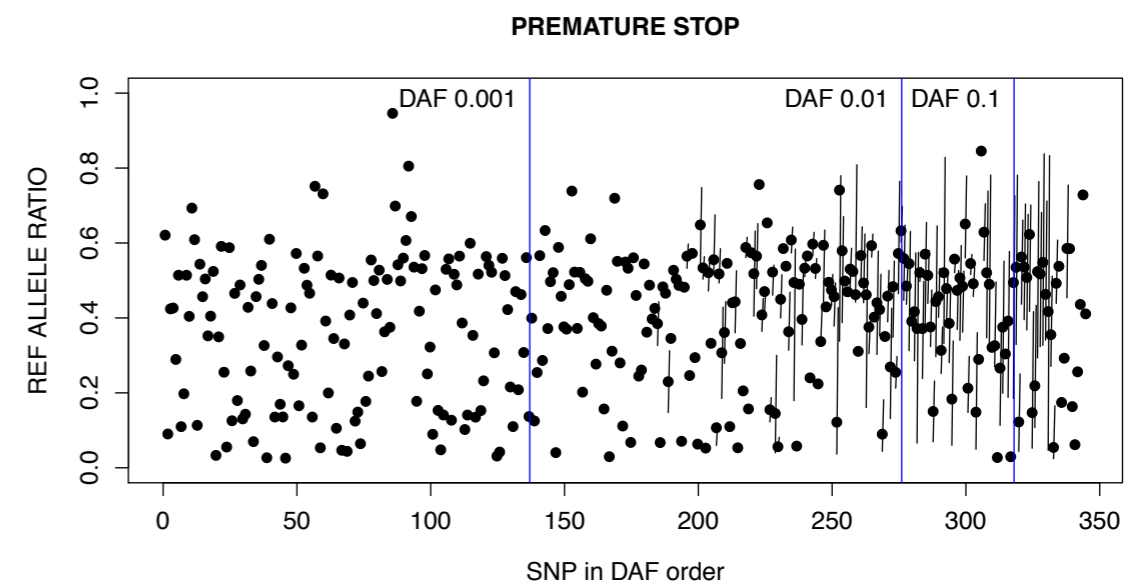
1. Add layer of Nonsense Mediated Decay

Predictions based on “50 bp termination code” rule.

2. Identify additional predictors of ASE/NMD signal:

- i. Relative position to start
- ii. Relative position to last splice junction
- iii. Protein Size or Number of Exons
- iv. Absolute Size of new transcript
- v. Conservation Score
- vi. Length of the Poly-A tail? (Is this measured)
- vii. Proximity of New/Original Stop to 3' UTR Start.
- viii. Additional predictors?

*Taken from Geuvadis BOG Poster.*



# Methods/Analysis

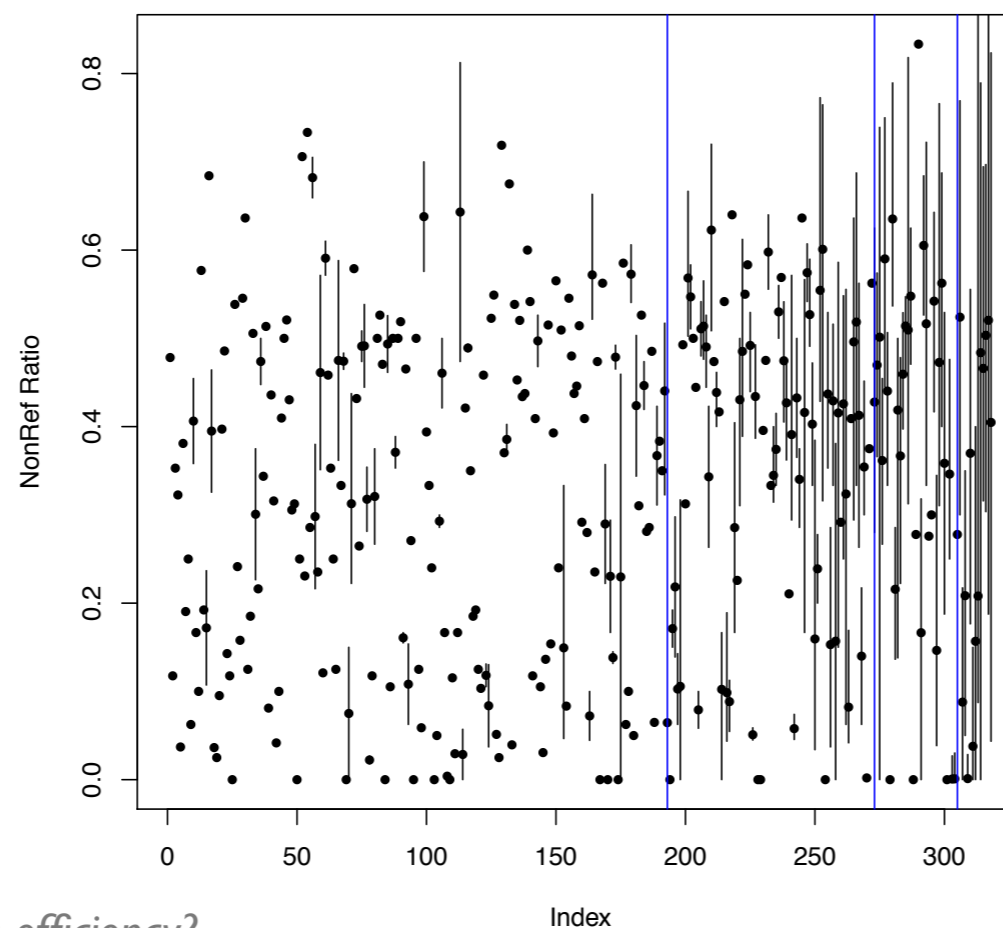
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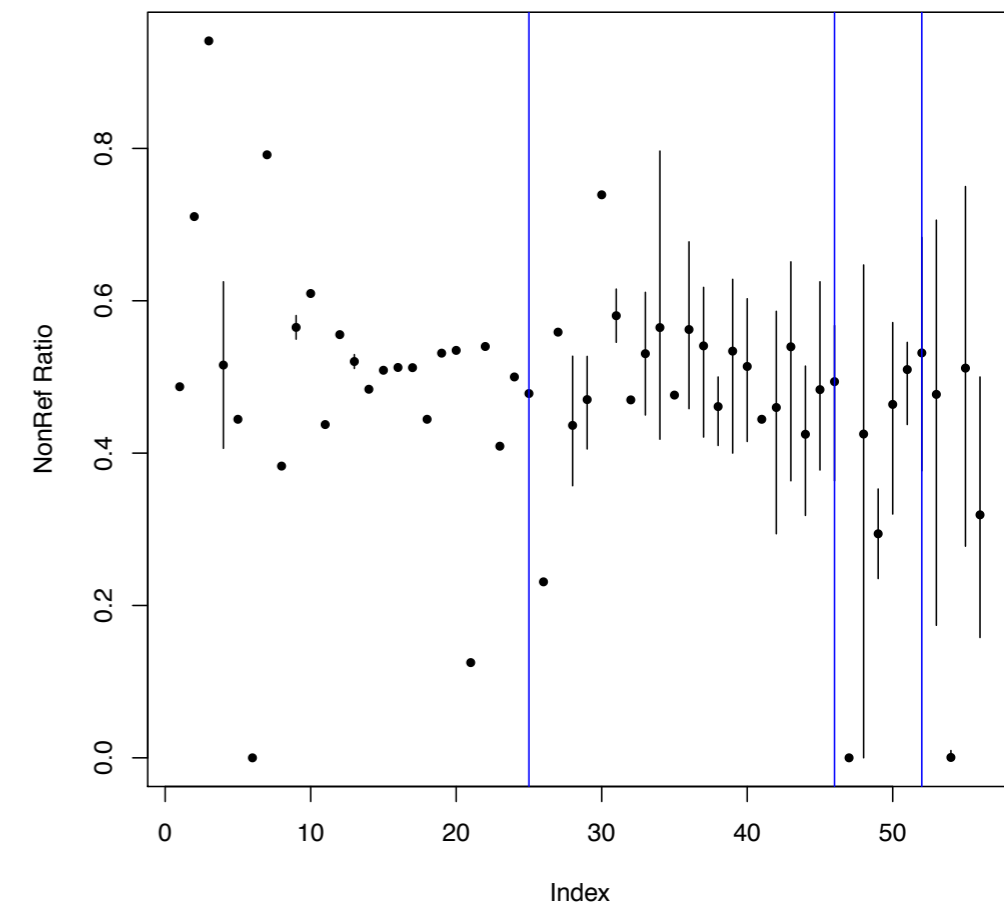
I. Add layer of Nonsense Mediated Decay

Predictions based on “50 bp termination code” rule.

*Predicted to Trigger NMD*



*Predicted to Escape NMD*



*Can we estimate NMD efficiency?*